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#### (57) Abstract

A fusion gene is provided comprising the cDNA of Renilla luciferase and the cDNA of the "humanized" Aequorea green fluorescent protein. The fusion gene was used to produce a novel protein, the "Renilla-GFP fusion protein", which displayed both the luciferase activity of Renilla luciferase, and the green fluorescence of GFP. The Renilla-GFP fusion gene is useful as a double marker for monitoring gene expression quantitatively in UV light and by enzyme activity.

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# RENILLA LUCIFERASE AND GREEN FLUORESCENT PROTEIN FUSION GENES CROSS-REFERENCE TO RELATED APPLICATIONS

The present Application is a International Application corresonding to United States Patent Application 08/771,850, filed December 23, 1996, entitled "The Construction and Expression of Renilla Luciferase and Green Fluorescent Protein Fusion Genes"; and is a Continuation-in-Part of United States Provisional Patent Application 60/027,657, filed October 4, 1996, entitled "The Construction and Expression of Renilla Luciferase and Green Fluorescent Fusion Genes in *E. coli* and Mammalian Cells," the contents of which are incorporated herein by reference in their entirety.

#### BACKGROUND

Green Fluorescent Protein (GFP) is a light emitting protein purified from the jellyfish Aequorea victoria. GFP can emit green light by accepting energy transfer from sources that include exogenous blue light and Renilla luciferase catalyzed reactions. The gene for GFP was cloned and its cDNA is a powerful reporter gene in a variety of living systems, including bacteria, fungi, and mammalian tissues. The UV light stimulated GFP fluorescence does not require cofactors and the gene product alone can be sufficient to allow detection of living cells under the light microscope.

By modifying the wild type GFP protein, red-shifted GFP variants with bright emission have also been produced. These variants include EGFP, GFPS65T and RSGF.

Recently, GFP was expressed in a human cell-line and in-vivo. C. Kaether, H.H. Gerdes.

Visualization of protein transport along the secretory pathway using green fluorescent protein.

FEBS-Lett. 1995, 369:267-71. "Humanized" GFP was synthesized with nucleotide changes that did not change the amino acid sequences with one exception.

Renilla luciferase is an enzyme purified from Renilla reniformis. The enzyme catalyzes the bxidative decarboxylation of coelenterazine in the presence of oxygen to produce blue light with an emission wavelength maximum of 478 nm? In Renilla reniformis cells, however, this reaction is shifted toward the green with a wavelength maximum of 510 nm due to an energy transfer to a Green-Fluorescent Protein.

The gene for Renilla luciferase (ruc) was cloned and its cDNA was shown to be useful as a reporter gene in various living systems D.C. Prasher V.K. Eckenrode, W.W. Ward, F.G. Prendergast, M.J. Cormier Primary structure of the Aequorea victoria green-fluorescent protein. Gene 1992 111:229-33. By providing appropriate promoters to the

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cDNA as gene cassettes, the gene was expressed in bacteria, transformed plant cells, and mammalian cells. The high efficiency of Renilla luciferase is a useful trait as a marker enzyme for gene expression studies.

Given the properties of GFP and Renilla luciferase, it would be useful to have a single protein combining the functions of both Renilla luciferase enzymes and GFP to monitor gene expression quantitatively by UV light excitation or qualitatively by enzyme activity measurements. Coulde in Englassia etc

### a on off and SUMMARY

According to one embodiment of the present invention, there are provided fusion gene constructs comprising the cDNA of Renilla luciferase and the cDNA of the "humanized" Aequorea green fluorescent protein. The fusion gene constructs were used to transform both prokaryotic and eukaryotic cells. One construct was expressed as a polypeptide having a molecular weight of about 65 kDa. This polypeptide, the "Renilla-GFP fusion protein," was bifunctional, displaying both the luciferase activity of Renilla luciferase and the green fluorescence of GFP The Renilla-GFP fusion gene is useful as a double marker for monitoring gene expression in living cells and quantitatively by enzymatic activity.

The invention includes a protein comprising a polypeptide having both luciferase and GFP activities, or biologically active variants of a polypeptide having both by luciferase and GFP, or a protein recognized by a monoclonal antibody having affinity to the 20 moopolypeptide having both luciferase and GFP activities. The polypeptide can be made by ad recombinant DNA methods and could a limit of the office of large to a

The invention further includes a high affinity monoclonal antibody that immunoreacts with the polypeptide. The antibody can have an Fc portion selected from the group consisting of the IgM class, the IgG class and the IgA class. The invention also includes a high affinity monoclonal antibody that immunoreacts with a polypeptide having both luciferase and GFP activities, quit sal sources don't or quillos tetrasaco no un sur suregi

The invention further includes a polynucleotide sequence coding for a polypeptide having both luciferase and GFP activities, or its complementary strands, and a polynucleotide sequence that hybridizes to such a sequence and that codes on expression for a polypeptide having both luciferase, and GFP, activities, or its complementary strands.

states of the invention further includes a purified and isolated DNA molecule comprising a polynucleotide coding for a polypeptide having both luciferase and GFP

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activities, or its complementary strands. The polynucleotide can comprise the sequence as set forth in SEQ ID NO: 1.

The invention further includes a vector containing a DNA molecule coding for a polypeptide having both luciferase and GFP activities. The polynucleotide can comprise the sequence as set forth in SEQ ID NO.1. The vector can be used to stably transform or and straight Vicinst florants through the size of a transiently transfect a host cell.

The invention further includes a method of making a polypeptide having both luciferase and GFP activities. The method comprises the steps of, first, culturing a microorganism transformed with a polynucleotide vector containing a gene cassette coding for a polypeptide having both luciferase and GFP activities. Next, the polypeptide having both 30 may . 2 g . luciferase and GFP activities is recovered.

The invention further includes a method of quantifying promoter activations and GFP fluorescence based on luciferase activity measurements. The method comprises the step of providing the polypeptide according to the present invention:

The invention further includes a method of making a monoclonal antibody that immunoreacts with a polypeptide having both luciferase and GFP activities. The method comprises the steps of, first, administering to a host a polypeptide having both luciferase and GFP activities in an amount sufficient to induce the production of antibodies to the polypeptide from the host's antibody-producing cells. Next, the antibody-producing cells are recovered from the host. Then, cell hybrids are formed by fusing the antibody-producing cell to cells 12 capable of substantially unlimited reproduction. Then, the hybrids are cultured. Next, the monoclonal antibodies are collected as a product of the hybrids

The invention further includes a method of monitoring gene expression. quantitatively and qualitatively in a cell using a gene fusion construct coding for a polypeptide 25 having both luciferase and GFP activities. The method comprises the steps of, first, providing a gene fusion construct coding for a polypeptide having both Renilla luciferase and GFP activity. Next, the gene fusion construct is introduced into the cell. Then, the cell containing the gene fusion construct is maintained in a manner alfowing the cell to express the polypeptide. Then, the cell is measured for luciferase and fluorescent activity. The construct can include a polynucleofide sequence as set forth in SEQ ID NO:124.000

The invention further includes a method of monitoring gene expression quantitatively and qualitatively in a cell using a gene fusion construct coding for a polypeptide

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having both luciferase and GFP activities. The method comprises the steps of, first, providing a gene fusion construct coding for a polypeptide having both luciferase and GFP activities. Next, the gene fusion construct is introduced into the cell. Then, the cell containing the gene fusion construct is maintained in a manner allowing the cell to express the polypeptide. Next, the cell is measured for luciferase and fluorescent activity.

#### FIGURES

These and other features, aspects and advantages of the present invention will become better understood with regard to the following description, appended claims, and accompanying figures where:

- Figure 1 is a schematic diagram showing the construction of a *Renilla* luciferase and "humanized" GFP fusion gene cassette according to the present invention for gene expression in *E. coli* where "RG," top, is the fusion gene cassette with the *Renilla* luciferase coding sequence (ruc) at the 5' terminus, and "GR," bottom, is the fusion gene cassette with the GFP coding sequence (gfp<sub>b</sub>) at the 5' terminus;
- Figure 2 is a schematic diagram showing the construction of Renilla luciferase and "humanized" GFP fusion gene cassette according to the present invention for gene expression in mammalian cells where "RG," top, is the fusion gene cassette with the Renilla luciferase coding sequence (ruc) at the 5' terminus, and "GR," bottom, is the fusion gene cassette with the GFP coding sequence (gfp<sub>b</sub>) at the 5' terminus;
- 20. Figure 3 is a map of the plasmids used for cloning and expression of the RG gene construct in E. coli (top) and the GR gene construct in E. coli (bottom);

Figure 4-is a map of the plasmids used for cloning and expression of the RG signed construct in mammalian systems (top) and the GR gene construct in mammalian systems (bottom);

- 25 Figure 5 are photomicrographs of cells transformed by the fusion genes using fluorescence microscopy and fluorescence imaging to show GFP activity:
  - Figure 6 are bar graphs of luciferase activity of the fusion gene constructs in E. coli (top) and mammalian cells (bottom);

30 a activity in E. coling a spectroscopic measurement of Renilla Inciferase activity and GFP

coli using anti-Renilla luciferase antibody; Antibody (1977) Antibody (1977) Color Williams (1977)

Figure 9 are photomicrographs of mouse embryonic stem cells using fluorescence image analysis demonstrating the expression of the RG fusion gene, and Figure 10 are photomicrographs of mouse embryos using fluorescence image analysis demonstrating the expression of the RG fusion gene.

DESCRIPTION

According to one embodiment of the present invention, there is provided a fusion gene comprising the cDNA of Renilta luciferase and the cDNA of the "humanized" Aequorea green fluorescent protein. According to another embodiment of the present invention, there is provided a single polypeptide that exhibits both Renilla luciferase and GFP activities. This bifunctional polypeptide can facilitate the identification of transformed cells at the single cell level, in cell cultures, transformed tissues and organs based on fluorescence of the polypeptide. At the same time, the polypeptide can also be used to quantify promoter activations and GFP fluorescence based on luciferase activity measurements.

The cDNA of Renilla rentformis luciferase (ruc) tias been cloned and used successfully as a marker gene in a variety of transgenic species. See, for example, Lorenz, W.W. McCann, R.O., Longiaru, M. and Cormier, M.J. Isolation and expression of a cDNA encoding Renilla rentformis luciferase. Proc. Natl. Acad. Sci. USA 1991; 88:4438-4442; Mayerhofer, R., Langridge, W.H.R., Cormier, M.J., and Szalay, A.A. Expression of, recombinant Renilla luciferase in transgenic plants results in high levels of light emission. The Plant Journal 1995, 7:1031-1038, and Lorenz, W.W., Cormier, M.J., O'Kane, D.J., Hua. D., Escher, A. A. Szalay, A.A. Expression of the Renilla reniformis luciferase gene in mammalian cells. J. Biolumin. Chemilumin. 1995, 11:31-37, incorporated herein by reference in their entirety. Similarly, the transfer and expression of Green-Fluorescent-Protein (GFP) cDNA from Aequorea victoria resulted in high levels of GFP in transformed cells that allowed convenient visualization of individual cells under the microscope. See, for example, Chalfie, M., Tu, Y., Euskirchen, G., Ward, W.W. and Prasher, D.C. Green fluorescent protein as a marker for gene expression. Science 1994; 263:802-805, incorporated herein by rimacod, ellisa quili am son i ès rais reference in its entirety.

The present invention involves the production of fusion genes from the cDNA of Renilla (ruc) and the cDNA of the "humanized" Aequorea GFP (gfp<sub>h</sub>). A description of "humanized" Aequorea GFP (gfp<sub>h</sub>) can be found, for example, in Zolotukhin, S., Potter, M., and Huaswirth, W.W., Guy, J., and Muzyczka, N. A "humanized" green fluorescent protein

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cDNA adapted for high-level expression in mammalian cells. J. Virology 1996: 70:4646-4654, incorporated herein by reference in its entirety.

The first fusion gene, designated the "RG fusion gene," SEQ ID NO 1 and shown at the top of Figures 1 and 2, contains the *Renilla* cDNA linked at the modified 3' end to a fifteen polynucleotide linker sequence encoding five amino acids, Ala-Ala-Ala-Ala-Thr, residues 312-316 of SEQ ID NO:1, followed by the 5' end of the intact GFP cDNA. The second fusion gene, designated the "GR fusion gene," SEQ ID NO:2 and shown at the bottom of Figures 1 and 2, contains the cDNA of GFP linked to a twenty-seven polynucleotide linker sequence encoding nine amino acids, Gly-Try-Gln-Ile-Glu-Phe-Ser-Leu-Lys, residues 239-247 of SEQ ID NO:2, followed by the 5' end of *Renilla* cDNA. Both genes were placed into prokaryotic pGEM-5zf(+) and eukaryotic pCEP4 expression vectors, and transformed into *E. coli*, and various mammalian cell lines, and microinjected into mouse embryos. PT<sub>7</sub> was the bacterial T7 promoter used for gene expression. P<sub>cmv</sub> was the CMV promoter used for gene expression in mouse fibroblast cells, embryonic stem cells and mouse embryos.

Unexpectedly, only cells transformed with the RG fusion gene gave strong fluorescence while the cells containing the GR fusion gene exhibited minimal response to UV light under the microscope. In contrast, luciferase measurements in homogenates of cells transformed with RG gene cassettes or with GR gene cassettes were indistinguishable from each other in both bacterial and mammalian cells. Further, spectrofluorimeter data indicated that there was energy transfer between Renilla luciferase and GFP in the RG fusion gene containing cells but did not indicate such energy transfer in cells containing the GR fusion gene. The protein expressed in the RG fusion gene containing cells was analyzed and found to be a 65 kDa polypeptide. A detailed description of the construction and expression of the fusion genes, and analyses of their protein products is given below.

Production of the Fusion Gene Constructs:

The vectors used for cloning and expression of the gene constructs in E. coli and mammalian systems were pGEM-5zf(+) (Promega) and pCEP4, respectively. Figure 3 is a map of the plasmids used for cloning and expression of the RG gene construct in E. coli, pGEM-5zf(+)-RG (top) and the map of the plasmids used for cloning and expression of the GR gene construct in E. coli, pGEM-5zf(+)-GR (bottom). Both were under the transcriptional control of T7 promoter. The E. coli strains which were transformed were DLT101 and DH5\alpha.

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Similarly, Figure 4 is a map of the plasmids used for cloning and expression of 1. ogy 1922the RG gene construct in mammalian systems, pCEP4-RG (top), and a map of the plasmids a conspirate di Ballogo alian used for cloning and expression of the GR gene construct in mammalian systems, pCEP4-GR (bottom). Both were under the transcriptional control of CMV promoter. The mammalian cell line that was transformed was LM-TK embryonic stem cells and embryos.

Five primers were designed for cloning the RG and GR gene constructs. Single underlines indicate Shine-Dalgarno sequences. Double underlines indicate the restriction sites. The start codons are in bold. Sequences in bold italics indicate the removal of stop codons abios has as said a from both ruc and gfp, genes.

Primer 1, SEQ ID NO:3:

RUC5: 5'CTGCAG (PstI)

AGGAGGAATTCAGCTTAAAGATG3'

ati a — A T — 19416: 19 . cmen Primer 2. SEQ ID NO:4: RUC3: 5'GCGGCCGC (Not I) TTG TTCATTTTTGAGAAC3'

Primer 3. SEQ ID NO:5:

GFP5:5 GGGGTACC (KpnI)

CCATGAGCAAGGGCGAGGAACT3

on was sala simmi soksa DA ed 15. Primer 4, SEQ ID NO:6:

dany bong ilagou aleo vico viberbequar GFP3: 5'GGGGTACC (Kpnl)

CCTTGTACAGCTCGTCCATGCCA3

ments in Language are in Felia Primer 5, SEQ ID NO:7:

GFP5a 5' CCCGGG (Smal)

### AGGAGGTACCCCATGAGCAAG3

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Renilla luciferase fusion gene (GR gene cassette) were constructed by removing the stop

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codons, and by adding restriction sites and Shine-Dalgarno sequences to the 5 end of the

products using PCR according to techniques known to those with skill in the art. The PCR

products were cloned using the pGEM-T system (Promega Corporation, Madison, WI). Primers were designed so that the downstream cDNA is in frame with the upstream cDNA.

The linker sequences are shown in Figures 1 and 2 and described above. After cloning, the RG and GR gene cassettes were under the transcriptional control of T7 in pGEM-5zf(+) vector and CMV in pCEP4 vector, which were used for expression in E. coli and mammalian vector and CMV in pCEP4 vector, which were used for expression in E. coli and mammalian vector and control of Section 2001 and to not sortice.

cells, respectively.

Determination of activity of fusion genes and their corresponding protein products:

GFP activity in vivo was visualized as follows. E. coli strain DH5α was न - Gir (bettern). Both were under the transformed with the plasmids pGEM-5zf(+)-RG and pGEM-5zf(+)-GR. Positive colonies were identified and cultured in LB medium with 100 μg/ml of ampicillin selection, according

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to techniques known to those with skill in the art. Twelve hours later, one drop of E. coli culture was put on a slide and visualized by fluorescent microscopy at 1000 x magnification. LM-TK cells were transfected with plasmids pCEP4-RG and pCEP4-GR using calcium phosphate methods known to those with skill in the art. The culture dishes were monitored using an inverted fluorescent microscope 12 hours after the transfection.

Luciferase activity was assayed as follows. An aliquot of transformed E. coli was used for a luciferase assay in a Turner TD 20e luminometer (Turner Designs, Sunnyvale, CA), both before and after IPTG induction. The results were recorded as relative light units. Mammalian cells harvested 36 hrs after transfection were measured for luciferase activity.

Corrected emission spectra were detected spectrofluorimetrically using a SPEX fluorolog spectrofluorimeter operated in the ratio mode. Fluorescence emission was excited at 390 nm. Bioluminescence emission was recorded with the excitation beam blocked following the addition of 0.1  $\mu$ g of coelenterazine in acidified methanol. Five spectra were averaged for each sample over a wavelength range from 400 to 600 nm.

The fusion proteins were isolated and immunoactivity detected as follows. 1 ml of E. coli. (OD<sub>600</sub>=1.0) was harvested. 400  $\mu$ l of cell suspension buffer (0.1M NaCl, 0.01 M Tris-HCl pH 7.6, 0.001 M EDTA, 100  $\mu$ g/ml PMSF) and 100  $\mu$ l of loading buffer (50 mM Tris-HCl pH 6.8, 2% SDS, 10% glycerol, 5% 2-mercaptoethanol) were added. The samples were boiled for 4 min and loaded to a 7.5%-20% gradient SDS-polyacrylamide gel.

20 Rolyclonal anti-Renilla luciferase was used as the primary antibody for detection and goat peroxidase anti-IgG (anti-rabbit) as the secondary antibody.

Referring now to Figure 5, there are shown photomicrographs of GFP activity in transformed E. coli cells (5A, left side) and LM-TK mouse fibroblast cells (5B, right side) by fluorescence microscopy and fluorescence imaging. As can be seen, individual E. coli cells and mammalian cells transformed with the RG fusion gene construct exhibited strong green fluorescence under oil immersion.

Referring now to Figure 6, there are shown bar graphs of luciferase activity of the gene constructs in E. coli (top) and mammalian cells (bottom). The white bars indicate activity before promoter induction. The black bars indicate activity after promoter induction. As can be seen, cells transformed with the RG fusion gene construct have significant luciferase activity, which is reduced 3-fold in the cells transformed with the GR fusion gene construct.

Referring now to Figure 7, there is shown a spectroscopic measurement of Renilla luciferase activity and GFP activity in E. coli transformed with various gene constructs. As can be seen, cells containing Renilla luciferase gene (short dashes) show only one emission peak at approximately 478 nm. Cells containing the GR gene fusion construct (light solid) also show one emission peak at approximately 478 nm, indicating Renilla luciferase activity only. By contrast, cells containing the RG gene fusion construct (heavy solid) show an emission peak at approximately 510 nm with excitation at 390 nm. Cells containing the RG gene fusion construct with the addition of coelanterizine (long dashes) show emission peaks at both approximately 478 nm and 510 nm indicating that the energy transfer between Renilla luciferase and GFP occurred in these cells. The lack of GFP activity in GR gene cassette transformed cell lines could be due to incorrect folding, due to the requirement for a free GFP C-terminus, or due to interference of the linker polypeptide with GFP activity in the fusion protein, among other possible explanations.

Referring now to Figure 8, there is shown a western blot used to detect fusion gene expression in E. coli using anti-Renilla luciferase antibody! Reading from left to right, the "C" lane shows the total protein extracted from non-transformed E. coli cells. The "R" lane shows the total protein extracted from E. coli cells transformed with the ruc gene alone. The "G" lane shows the total protein extracted from E. coli cells transformed with the gfp, gene alone. The "RG" lane shows the total protein extracted from E. coli cells transformed with the RG fusion gene cassette. The "GR" lane shows the total protein extracted from E. coli cells transformed with the RG fusion gene cassette.

As can be seen, protein extracted from E. coli cells transformed with the ruc gene alone produced a band with a molecular weight of about 34 kDa. Protein extracted from E. coli cells transformed with the RG fusion gene cassette produced a band with a molecular weight of about 65 kDa. Protein extracted from E. coli cells transformed with the GR fusion gene cassette produced a band with a molecular weight of about 34 kDa. These data imply that cells transformed with the GR fusion gene cassette produced luciferase but did not produce cells transformed with the GR fusion protein production by cells transformed with the GR fusion protein production by cells transformed with the GR fusion protein production by cells transformed with the GR

Referring now to Figure 9, there are silown photomicrographs using the same of the RG fusion gene in mouse

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embryonic stem cells transformed by electroporation procedures. Transformed colonies were selected based on GFP activity under fluorescence microscopy.

Referring now to Figure 10, there are shown photomicrographs using fluorescence image analysis demonstrating the expression of the RG fusion genes in mouse embryos. The embryos were injected with the linearized RG plasmid, and *in vitro* cultured. The expression of GFP activity was monitored daily by fluorescent microscope and recorded by an imaging collection system.

Based on this data, we conclude that the RG fusion construct disclosed herein can be expressed in both prokaryotic and eukaryotic cells to produce a bifunctional polypeptide that exhibits both *Renilla* luciferase and GFP activity. This bifunctional polypeptide is a useful tool for identification of transformed cells at the single cell level based on fluorescence. It allows the simultaneous quantification of promoter activation in transformed tissues and transgenic organisms by measuring luciferase activity. The dual function of this protein allows the monitoring of bacterial cells in their living hosts and the differentiation of cells in the developing embryo and throughout the entire animal.

Although the present invention has been discussed in considerable detail with reference to certain preferred embodiments, other embodiments are possible. Therefore, the spirit and scope of the appended claims should not be limited to the description of preferred embodiments contained herein.

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	ATG	ACT	TCG	AAA	GTT	TAT	GAT	CCA	GAA	CAA	AGG	AAA	CGG	ATG	ATA	ACT Thr		48
	Met	Thr	Ser	Lys	Val	Tyr	Asp	Pro	GIU	10	Arg	rys	Arg	1100	15		•	
								·.				7 7 M		CTT.	CD.TT	TCA		96
	GGT	CCG	CAG	TGG	TGG	GCC	AGA	TGT	LVS	.Gln	Met	Asn	GTT Val	Ten	Asp	Ser		
	GIĀ	Pro	GIII	20	LP	7.10		- 1 -	25	•				30				
•				mam	·mvm	C D T	TCA	GAD.	444	CAT	GCA	GAA	AAT	GCT	GTT	АŢТ		144
•	Phe	ATT	AAT	Tyr	Tyr	Asp	Ser	Glu	Lys	His	Ala	Glu	ASII	Ala	Val	Ile	*	
•			35		- -	٠		4,0					45	*				
	TTT	TTA	CAT	GGT	AAC	,GCG	GCC	TCT	TCT	TAT	TTA	TGG	CGA	CAT	GTT	GTG		192
	Phe	Leu	His	Gly	Asn	Ala	Ala	Ser	Ser	Tyr	.Fe <i>i</i> r	Trp 60	Arg	His	v.a1	vai		
		50		1			55					-						240
	CCA	CAT	ATT	GAG	CCA	GTA	GCG	CGG	TGT	ATT	ATA	CCA	GAT	CTT	ATT Ile	GGT Glv	-	240
	Pro 65	His	Ile	Glu	Pro.	70 70	ATG	Arg	Cys	TIE	75	110	Asp			80	•	٠.
							m		20.20.00	~ ~m	T CT	ייי צויירי ייי צויירי	שכה	מידים	СТТ	GAT	•	288
	ATG	GGC	AAA	TCA	GGC Glv	AAA 'Lv.s	Ser.	GLY	AAT	Gly	Ser	Tyr	AGG Arg	Leu	D,C G	تو ت		
	.1100	O = 3	-,-			-				0.0					.95			

CÁT His	TAC Tyr	AAA Lys	TAT Tyr 100	CTT Leu	ACT.	GCA: Ala	TGG <sub>1</sub>	TTT Phe 105	GAA Glu	CTT Leu	CTT Leu	AAT Asn	TTA Leu 110	CCA. Pro	AAG Lys	Ι: ε	3,36
AAG Lys	ATC Ile	AAT Ile 115	TTT Phę	GTC Val	GGC Gly.	CAT:	GAT: Asp 120	TGG. Trp	GGT. Gly	GCT Ala	TGT- Cys	TTG Leu 125	GCA Ala	TTŢ. Phe	CAT.		3,84
TAT Tyr	AGC Ser 130	TAT Tyr	GAG Glu	CAT. His	CAA Gln	GAT Asp 135	AAG Lys	ATC Ile.	AAA Lyş	GCA Ala	ATA Ile 140	GTT Val	CAC His	GCT Ala	GAA Glu	 :	432
AGT Ser 145	GTA Val	GTA Val	GAT.	GTG: Val.	ATT. Ile 150	GAA.	TCA Ser	TGG: Trp	GAT:	GAA Glu 155	TGG Trp	CCT Pro	GAT Asp	ATT Ile	GAA Glu 160	•	480 ·.
GAA Glu	GAT Asp	ATT Ile	GCG Ala	TTG: Leu: 165:	Tle	AAA Lys	TCT. Ser	GAA Glu	GAA Glu 170	Gly	GAA Glu	AAA Lys	ATG Met	GTT Val 175	TTG Leu	. 5	528
GAG Glu	AAT Asn	AAC Asn	TTC Phe 180	TTC Phe	GTG Val	GAA Glu	ACC Thr	ATG Met 185	TTG	CCA Pro	TCA Ser	AAA Lys	ATC Ile 190	ATG Met	AGA Arg		576
AAG Lys	TTA Leu	GAA Glu 195	CCA Pro	GAA Glu	GAA' Glu	TTT.	GCA Ala 200	GCA: Ala	TAT Tyr	CTT: Leu	<u> </u> Gl <sub>i</sub> u	CCA Pro 205	TTC Phe	-AAA Lys	GAG Glu		624
AAA Lys	GGT Gly 210	GAA Glü ∂3	GTT Val	CGT	CGT.	CCA Prod 215	TACA:	Leu.	TCA Ser	TGG Trp	CCT. IPro. 220	CGT Arg	GAA, Glu	AŢÇ Iļę	CCG Pro		6 <b>72</b>
TTA Leu 225	GTA Val	AÄÄ Lÿs	GGT Gly	GGT Gly	AAA Lys 230	Pro	GAC Asp	GTT Val	GTA Val	CAA Gln 235	AŢŢ	GTT: Val:	AGG Arg	AAT, Asn	TAT Tyr 240		. 720 
AAT Asn	GCT Ala	TAT Tyr	CTA	CGT Arg 245	Ala	AGT Ser	GAT: Asp	GAT: Asp	TTA: Leua 250	Pro	AAA Lys	ATG: Met:	TTT Phe	ATT Ile 255	GAA Glu	2-7 173 11	7.68 Q 181
TÉĞ Ser	GAT Asp	CĈĀ: PĒŌ:	GGA G1 y- 260	TTC: Phe	TTT Phe	TCC. Ser	TAA Asn	GCT: Ala: 265	ATT [11e	gGTT Val∵	; GAA: ⊝GLu	CTA CCC	GC.G- Al.a. 270	AAG Lys	AAG Lys	DAĄ J	816
-ŤŤŤ Phe	CCT Pro	AAT Asn- 275	ACT. The	GAA Glu	TIT Phe	Va.1.	Lys 280	GTA Val:	AAA Lys_	GGT Gly	Leu	CAT His 285	TŢŢ.	TCG Ser	CAA Gln		864
GAA Glu	GAT Asp 290	GCA Ala	CCT Pro	GAT Asp	GAA Glu	ATG Met 295	GGA: Gly	AAA: Ly.s:	TAT.	ATC Ile	Lys	TCG Ser	ŢŦÇ. Phe	GTT Val	GДu GДu	7 1. 7 (p	912 
CGA Arg 305	GTT Val	CTC Leu	AAA Lys	AAT Asn	GAA Glu 310	CAA Gln	GCG Ala	GCC Ala	GCC Ala	Ala :3:15	Thr	Met	Ser	Lys	Gly -320	*	960
GAG Glu	GAA Glu	CTG Leu	TTC Phe	ACT Thr 325	GGC Gly	GTG Val	GTC Val	Pro	11e ∵330	aCTC aLeu ca	GTG	GAA: Glu: L::IV	erica ienen ienen	GAT Asp 335	GGC		1008
GAT Asp	GTG Val	Asn	Gly 340	His	Lys	Phe	Ser	Val 345	AGC Ser	GGA Gly	TGAG Glu	GGT Gly	GAA Glu 350	G1 y	GAT Asp	: ::::::::::::::::::::::::::::::::::::	

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Ala	Thr	Tyr-(	Gly :	Lys	Leu	The	160 360	ъys-	Phe	, 116	Cys.	365	THES	<b>G</b>	-	. "	ine a is t
CTC Leu	CCT Pro 370	GTG (	CCA Pro	rgg Trp	Pro	ACA Thr 375	CTG Leu	GTC Val	ACT Thr	ACC: The	TTC' Phe 380	ACC Thr.	TAT Tyr	GGC Gly	GTG Val::	~ m,4,1	152
CAG Gln 385	TGC Cys	TTT Phe	TCC . Ser	AGA Arg	TAC Tyr 390	CCA Pro	GAC Asp	CAT His	ATG Met	AAG Lys 395	CAG Gln	CAT-	GAC Asp:	.TTT.= .Phe	TTC T Phe T 400	) ( )	1200
AAG Lys	AGC Ser	GCC Ala	Met <sup>.</sup>	CCC Pro 405	GAG Glu	GGC G1 y	TAT Tyr	GTG Val	CAG Gln 410	GAĞ. Glu	AGA Arg	ACC' Thr	ATC Ile	TTT Phe 415	TTC Phe	3	1248
AAA Lys	GAT Asp	GAC Asp	GG <b>G</b> G1 y 420	AAC Asn	TAC Tyr.	AAG Lys	ACC Thr	CGC Arg 425	GCT: Ala	GAA Glu	GTC Val	AAG Lys	TTC Phe 430	GIU.	GGT Gly		1296
GAC Asp	ACC Thr	CTG Leu 435	GTG Val	AAT NaA	AGA Arg	ATC Ile	GAG Glu 440	Leu	AAG Lys	GGC Gly	ATT Ile	GAC Asp 445	TTT	AAG Lys	GAG Glu	٠.	1344
Asp	Gly 450	Asn	Ile	Lèü	G1 y	His 455	Lys	_Leu	Glu	Tyr	460	Tyr.	ASII	.,ser,	CAC	• •	1392
Asn 465	Val	Tyr	Ile	Met	Ala 470	Asp	:Lÿ:s :	GIn	Lys	.:ASn:	.GI y.	ite	.uya	, √A € ₩'	400		1440
Phe	Lys	Ile	Arg	His 485	Asn	Ile	Gľú .:	Asp	490	Ser	vaı	GEN.	Leu	495	gGAC <sub>C</sub> , (Asp		1488
His	туг	Gl n	Gln 500	Ash	ThE	Pro	lle	505	Asp	GIY	PEO	· .val	510	y Y	1144		
Asc	Asn	His 515	Tyr	Leu	`Ser	Thr	520	ser	: Ala	Leu	ser	525	) (ASP	FIU		1	
GAP Glu	A AAG Lys 530	Arg	GAĆ Asp	CAC His	Met	. va.1	CTO	ı Let	1. GT.	TTT Phe	· · val	1 11.1	: GCI	GCT Ala	i, ege id ege	. 😅	1632 s. 1
Ile	C ACA E Thi	A CAT	GGG G1y	ÁTÓ Met	. Asp	GA(C)	ı. Lei	TAC Ty	E AAC	TGA		A + 13 N + 2 Z	ja is k Diliya S	B (1) 2 (2);		 1211 125	1665
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120			SEQU	JEŇC	E DES	SCRI	PTIO	N : . S. .: ∋	E. 1	V 120		1 4	2		TAX	1000	48
AT Me	t Se	r Lys	s Gly	GA G1	u Gl	u Le	u Ph	e Th	r Gl	y Va.	l Va	1 Pr	יבביט	e Le 1	a var		

WO 98/1460	5			PCT/US97/1	7162
GAA CTG GAT Glu Leu Asp					96

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GGT Gly	GAA Glu	GGT GAT Gly Asp 35	GCC Ala	ACA Thr	TAC Tyr	GCA Gly 40	AAG Lys	CTC Lėù	ACC Thr	CTG Leu	AAA Lys 45	TTC Phe	ATC Ile	TGC Cys	. **.	144
ACC	ACT Thr 50	GGA AAG Gly Lys	CTC Leu	CCT Pro	GTG Val 55	CCA Pro	TGG Trp	CCA Pro	ACA Thr	CTG Leu 60	GTC Val	ACT Thr	ACC Thr	TTC Phe	· .	192
ACC Thr 65	Tyr	GGC GTG Gly Val	CAG Gln	TGC Cys 70	TTT	TCC Ser	AGA Arg	TAC Tyr	CCA Pro 75	GAC Asp	CAT His	ATG Met	AAG Lys	CAG Gln 80	· · · ·	240

CAT GAC TTT His Asp Phe	TTC AAG AGG GGC Phe Lys Ser Ala 85	ATG CCC GAG GGC Met Pro Glu Gly 90	TAT GTG CAG GAG Tyr Val Gln Glu 95	AGA 288 Arg
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ACC	מתכ	ጥጥጥ	ጥጥር	מממ	CAT	GAC	GGG	ממ	<b>ጥ</b> አ ር	חחר	. N.C.C.		C C TT	C D D	CDC	336
Thr	Tle	Phe.	Phe	1.75	ASD	Asn	Glv	Den	Tyr	Tue	24CC	7560	חום	CAN	Val	336
		,	100	LyJ	,,,	طوين	O <sub>z</sub> y	105				Ard	•		.var	· ·
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AAG TTC Lys Phe	GAA: GGT	GAC ACC	CTG	GTG I	AAT	AGA	ATC	GAG	CTG	AAG	GGC -	ATT	-	384
Lys File	GIU GIY	wah int	nie u	Val. 1	ASII	Arg	$_{11e}$	ين⊥ س	.reu	,Lys	GLY .	ile -		
	115		."	120					125				- '	

GAC	TTT	AÁĞ	GÁĞ	GÁT	GĠA	AAC	ATT	CTC	GGC	::CAC	AAG	(CTG	GAA	TAC	AAC (+ .)	:	.432
Asp	Phe	Lys	Glu	Asp	Gly	Asn	Ile	Leu	Gly	His	Lys	: Leu	Glu	Tyr	Asn		
	130					135	54				140			•		- :	

TAT AAC	rec các áat	GTG TAC	ATC ATG GCC	GAC AAG	CAA"AAG (AATGGC," t	480
Tyr Asn S	Ser Hrs Asn	Val Tyr	Ile Met Ala	Asp Lys	Gln LyscAsn-Gly	
145		150	۾ (	155	160	

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DTC	DDG	CTC	חתת	TTC	DAG	`DTC	DCD	CAC	カカケ	J. June	CDC	~ n m	CCD 'MCC	. cmc i.	528 زخیر
71.0	7070	6, 0	777	110	منح	AI C	AGA	CAC	AAC	WIT I	GAG	GA 1	GGA ILC		JUL 028
TIA	LUC	Va'l	D C D	Dhip	1410	T10	Dra	. Hire	"N'en	T 3	Jen 1	n	Gly:Ser	57°- 1	
110	23	V CL	7.311	1110	цyэ	116	~ry	1112	VOII	. TIE.	-GT II	· Asp.	Gry-Ser	val	
				165					.:1.7 N				3175		

CAG	CTG	GCC	GAC	CAT TAT	CAA CAG	AAC	ÄČT CCA	ATC	ĠĠĊ-ĠĀ	୍ର ଓଡ଼ିଆ	CCT		- 5	76
Gln	Leu	Ala	Ásp	His Tyr	Gln Gln	Asn	ThrePro	Tie	GIV As	Glv	Prom:	9.	_	
			180	~ * t*	_	185			19	D				

GTG	CTC	CTC	CCA	GAC	AAC	CAT	TAC	CTG	TCC	ACC	CAG	TCT	GCC CTG	TCT	 624
Val	Leu	Leu	Pro	Asp	Asn	His	Tyr	Leu	'Ser	Thr	Ġĺn	Ser-	Ala Leu	Ser-	 :
		195				. **	200			-	: ::	205		J 161	

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AAA	GAT	ĊCČ	AAC	GAA	AAG	AGA	GAC	"ĈÃC	ATG	GTC	CTG	CTG	GAG TTT GT	G 4		672
1.05	Asn	Pro	Asn	Gin	Tive	Dra	Adn	Hie	Mat'	175.1	1000	7-01	Glu Phe Va		S 22 77	0,2
2,2		110	7311	GIG	- y - y - y			1113	MEC				Giu Phe va	1		
•	210					215	•				220	* 2				

ACC GC Thr Al 225	T GCT a Ala	GGG G1 y	ATC .	ACA CAT Thr His 230	GGC ATG GAC Gly Met Asp	GAG CTG Glu Leu 235	TAC AAG G	G TAC Ly Tyr	ે કે કે આ કે કે આ કે	720
				230		233	•	240		

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CAG	ATC	GAA	TTC	· AGC	TTA	AAG	ATG	ACT	rcci	AAA	CTT.	ידעידי	CAT	CCA.	GAĀ Glu	760
				-,			4.			,,,,,		****	GAL		GAA.	 100
Gln	Ile	G1u	Phe	Ser	Leu	Lvs	Met	ጥኮሎ∴∢	Ser	LVS	Val	Tive	ີກເກັ	70.70	C1	
						-,-			75-	Lys	V 44 1	- y -	MSP	ELU	Giu	
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CAA	agg aaa	CGG	ATG ATA	ACT GGT	CCG CAC	TGG TGG	GCC AG	TGT AAA		816
Gln i	Arg Lys	Arg	Met : Ile	Thr Gly	Pro Glr	Tro Tri	o Ala Ard	Cys Lys	× 5	
	•	260		•	265		270	)		

		the state of the s		• • •
CAA ATG AAT GTT	CTT GAT TCA TTT	ATT AAT TAT TAT	GAT TCA GAA AAA	864
Gln Met Asn Val	Leu Asp Ser Phe	Ile Asn Tur Tur	Asp Ser Glu Tue	3 3

				25.5 T (4.5										1920 4	0.19 0.19			441. 1441.
			275					280		. :			285		Ç			
	CAT His	GCA Ala 290	GAA Glu	AAT Asn	GCT Ala	ĠŤŤ Val	ATT Ile 295	TTT Phe	TTA Leü	CAT His	GGT Gly	AAC Asn 300	GCG Ala	GCC Ala	TCT Ser	TCT Ser	Subjective Control	912
•	TAT Tyr 305	TTA Leu	TGG Trp	CGA Arg	CAT	GTT Val 310	GTG Val	CCA Pro	CAT	ATT Ile	GAG Glu 315	CCA Pro	GTA Val	GCG Al'a	cgg Arg	TGT Cys 320		960
	ATT Ile	ATA Ile	CCA Pro	GAT Asp	CTT Leu 325	ATT Ile	GGT Gly	ATG Met	GGÇ G1 y	AAA Lys 330	TCA Ser	GGC Gly	AĀĀ Lys	TCT Ser	GGT Gly 335	AAT Asn	;	1008
3 :	GGT Gly	TCT Ser	TAT	AGG Arg 340	TTA Leu	CTT Leu	GAT Asp	CAT His	TAC Tyr 345	AAA Lys	TAŤ Týr	CTT Leu	ÀCT Thr	GCA Ala 350	TGG Trp	TTT' Phe	· · · · · · · · · · · · · · · · · · ·	1056
	GAA Glu	CTT Leu	CTT Leu 355	Asn	TTA Leu	-CCA Pro	AAG Lys	AAG Lys 360	ATC Ile	ATT Ile	TTT Phe	GTC Val	©GC Gly 365	CAT	GAT Asp	TGG. Trp		1104
	GGT Gly	GCT Ala 370	Cys	ŤTG ′Leu	GCA -Ala	TTT Phe	CAT His 375	ТАТ Туг	AGC Ser	TAT	GAG Glu	CAT His 380	CAA GIn	GAT Asp	AAG Lys	ATC Ile		1152
3.4	AAA Lys 385	Ala	ATA Ile	GTT Val	-CAC JHis	GCT Ala 390	∴G1′u	AGT Se'r	GTA Val	GTA Val	Asp	GTG Val 5	ATT	GAA Glu	TCA Ser	TGG. Trp 400		1200
: · 3	GAT Asp	GAF Glu	∴TGG i Trp	GCT	GAT Asp 405	Ile	GAA ∍Glu	GAA Glu	-GAT Asp	ATT Ile 410	GCG Ala	T/TG Leu	ATC Ile	AAA Lys	TCT Ser 415	GAA Glu	÷ 9	1248
	GAA Glu	GGZ Gly	/: Glu	LAAA Lys 3420	Met	∵GTT ∖Val	TŢG ⊹ţeu	_GAG _Glu	AAT ASN 425	Asn	TTC Phe	TTC Phe	GTG Val	GAA Glu 430	THE	ATG Met		1296
10	TTC Leu	CC2	TCF Ser 435	Lys	rATC dille	ATG Met	-AGA Arg	AAG Lys 440	Leu	-GAA Glu	CCA Pro	GAA Glu	GAA Glu 445	hué	GCA Ala	GCA Alà	· 773.77 9.7	1344
У С	TAT	Le	GA/ u Glu 0	A, ÇÇP 1. Pro	Phe	AAA Lys	GJ u	ı Lys	- GGT - Gly	GAA Glu	Val	Arg 460	Arg	CCA Pro	ACA Thr	TTA Leu		1392
	5 TC/ Se: 46!	Tr	g cc	o Arc	GA Glu	$_{i,z}$ lle	CCG Pro	TTA	GTA Val	Lys	GGT Gly 475	٧.٠٤ ب	Lys	CCT Pro	GAC Asp	var	30 438	1440
~ .		1 G1	n Th	e Val	48:	j Asr	Ţŷı	: Aşr	Y Ala	490	Let )	Arg	Ala F	i ser	495	, wsb	3	1488
		u Pr	о ту	5 Me 3500	); -:	= , <del>1</del> ,1		5 7.1	50	5		1		3510 122	) # 1.7	, ¥ . î		1536
÷.	11	e Va	177G1 3751	A GGO u Gl	ÿ A1	a Ly	s Ly:	520	D: - :	orası oli 330	1 1n:	GET.	⇒52:	5 TeM	n iA Ain Sec			1584
:	AA Ly	- ~ 1	- i. The	. 2. 11 -	~~ Dh.	<b>A</b>	77 . [ - 1 ]	n. :-!		r GCA	4 . P.	JASI	J - G I	u - 1-1-	G GG/	AAA y Lys	(4) <sup>7</sup> 1 21	1632

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TAT ATC AAA TCG TTC GTT GAG CGA GTT CTC AAA AAT GAA CAA TAA 1677
                      Tyr. Ile Lys. Ser Rhe Val Glu Arg Val Leu Lys Asn Glu Gln *** 545
                                                                                                                                                                                                                                            我 医聚乙基 人名 经
                              (3) INFORMATION FOR SEQ ID NO:3:
                                                           (1) SEQUENCE CHARACTERISTICS:
                                                                                  (A) LENGTH: 29 base pairs
                                                                                  (B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear
 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
                      CTGCAGAGGA GGAATTCAGC TTAAAGATG
                             (4) INFORMATION FOR SEQ ID NO:4:
                                                          (1) SEQUENCE CHARACTERISTICS:
                                                                                        (A) LENGTH: 26 base pairs
                                                                                                                                                                                                                                               - 11 11 1 1 1 1 1 T
                                                                                        (B) TYPE: nucleic acid
                                                                                                                                                                                                                                         A Committee of the Comm
                                                                                        (C) STRANDEDNESS: single
                                                                                        (D) TOPOLOGY: linear
                                                          (x1) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
                           GCGGCCGCTT GTTCATTTTT GAGAAC
                             (5) INFORMATION FOR SEQ ID NO:55 ( WARRED AN ELECTION ALBERT COLD
                                           (1) SEQUENCE CHARACTERISTICS:
                                                                              (A) LENGTH; 30 base pairs 10 god as story a Akit t
                                                                                (B) TYPE: nucleic acid
                                                                                       (C) STRANDEDNESS: single
                                                                                       (D) TOPOLOGY: linear
      West of the (wife sequence: Description: sequ
               GGGGTACCCC ATGACCAACGCCQQAGGAACTC CONTRACTOR CONTRACTOR
                         (6) INFORMATION FOR SEQ ID NO:6:
                                                                                                                                                                                                                                                                               en element of the element
                                                      (1) SEQUENCE CHARACTERISTICS:
                and of the first of the dength: 36 base pairs for the local plants 1941 A 191
                                                                                       (B) TYPE: nucleic acid
                                                                                       (C) STRANDEDNESS: single Latinus 430 par agarefrout that parend
                                                                                       (D) TOPOLOGY: linear
      THE CHARLES SEQUENCE DESCRIPTION SEQUENCES & Las De. John A. . I .
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#### WE CLAIM:

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1. A protein comprising a polypeptide having both luciferase and GFP activities or biologically active variants thereof.

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- 2. A recombinant protein according to claim 1.
- 3. A protein according to claim 1, having an amino acid sequence as set forth in SEQ ID NO:1.
- 4. A high affinity monoclonal antibody which immunoreacts with the polypeptide of claim 1.
- 5. The antibody of claim 4 having an Fc portion selected from the group consisting of the IgM class, the IgG class and the IgA class.
- 6. A protein recognized by a monoclonal antibody having affinity to the polypeptide of claim 1.
  - 7. The protein of claim 1 in purified and isolated form.
- 8. A DNA sequence coding for a protein according to claim 1, or its complementary strands.
- 9. A DNA sequence which hybridizes to a DNA sequence according to claim 8 and which codes on expression for a polypeptide having both luciferase and GFP activities, or its complementary strands.
- 10. A high affinity monoclonal antibody which immunoreacts with a polypeptide having both luciferase and GFP activities.
- 11. A purified and isolated DNA molecule comprising a polynucleotide coding for a polypeptide having both luciferase and GFP activities, or its complementary strands.
- 12. The DNA of claim 11, wherein the polynucleotide comprises the sequence as set forth in SEQ ID NO:1.
- 13. A vector containing a DNA molecule coding-for a polypeptide having both luciferase and GFP activities.
- 14. The vector of claim 13, wherein the polynucleotide comprises the sequence as set forth in SEQ ID NO:1.
- 15. A prokaryotic or eukaryotic host cell stably transformed or transfected by the vector of claim 13.
  - 16. A method of making a polypeptide having both luciferase and GFP activities, the method comprising the steps of:

- polypeptide having both luciferase and GFP activities; and
  - (b) recovering the polypeptide having both luciferase and GFP activities.
- 17. A method of quantifying promoter activations and GFP fluorescence based on luciferase activity measurements, the method comprising the step of providing the polypeptide according to claim 1.
- 18. A method of making a monoclonal antibody which immunoreacts with a polypeptide having both luciferase and GFP activities, the method comprising the steps of:
  - (a) administering to a host a polypeptide having both luciferase and GFP activities in an amount sufficient to induce the production of antibodies to the polypeptide;
    - (b) recovering the antibody-producing cells from the host;
  - (c) forming cell hybrids by fusing the antibody-producing cell to cells capable of substantially unlimited reproduction;
    - (d) culturing the hybrids; and
    - (e) collecting the monoclonal antibodies as a product of the hybrids.
- 19. A method of monitoring gene expression quantitatively and qualitatively in a cell using a gene fusion construct coding for a polypeptide having both luciferase and GFP activities, the method comprising the steps of:
  - (a) providing a gene fusion construct coding for a polypeptide having both Renilla luciferase and GFP activity;
    - (b) introducing the gene fusion construct into the cell:
  - (c) maintaining the cell containing the gene fusion construct in a manner allowing the cell to express the polypeptide; and
    - (d) measuring the cell for luciferase and fluorescent activity.
- 20. The method of claim 19, where the step of providing comprises providing a construct including a polynucleotide sequence as set forth in SEQ ID NO:1.
- 21. A method of monitoring gene expression quantitatively and qualitatively in a cell using a gene fusion construct coding for a polypeptide having both luciferase and GFP activities, the method comprising the steps of:
  - (a) providing a gene fusion construct comprising the protein of claim 1;
  - (b) introducing the gene fusion construct into the cell;

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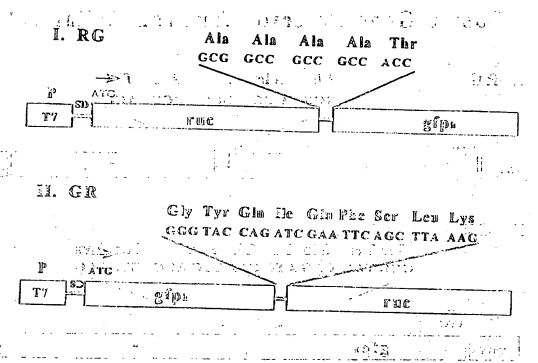
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FIG. 1

### Fusion Gene Cassettes for E. coli

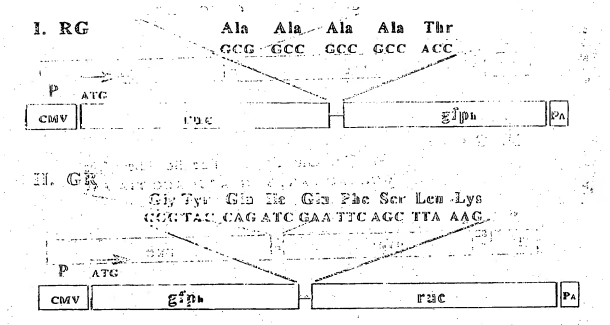


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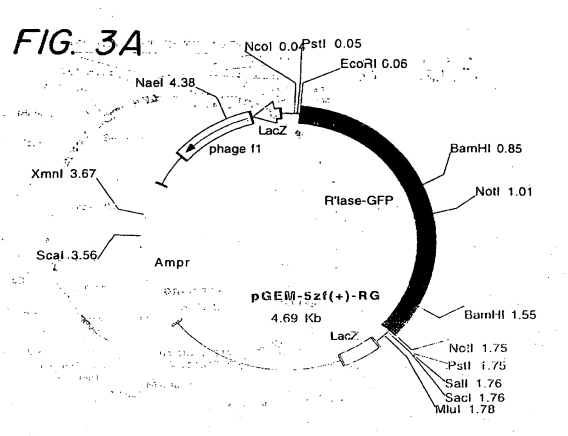
FIG. 2

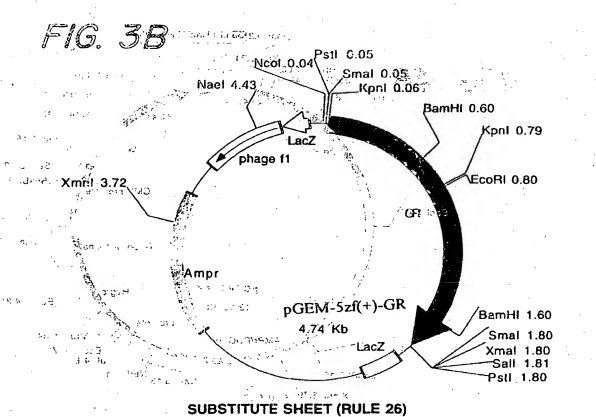
### Fusion Gene Cassettes for Mammalian cells



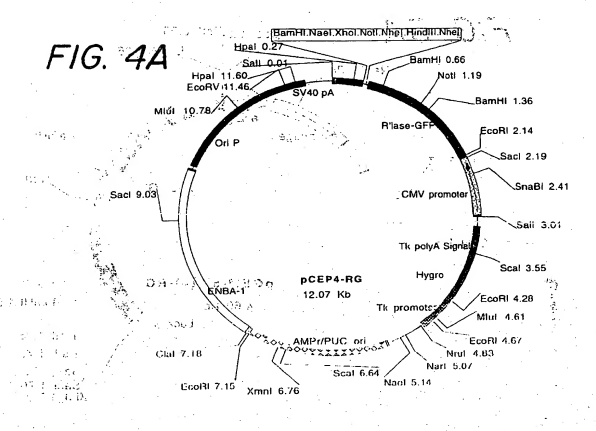
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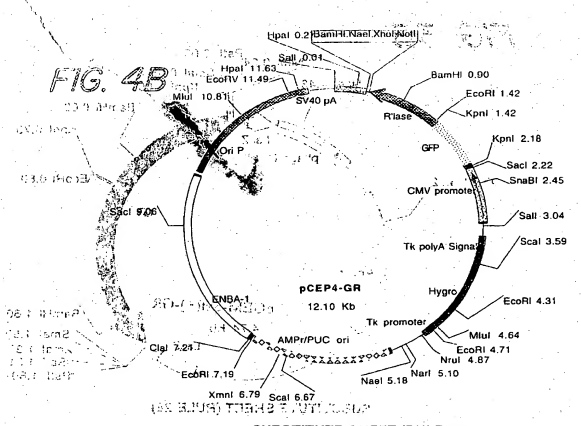
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FIG. 5A

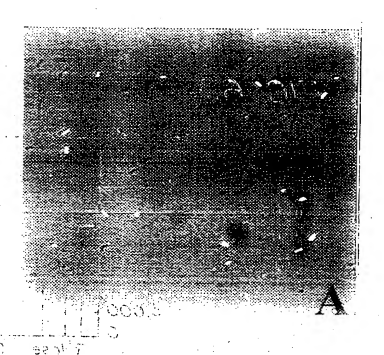


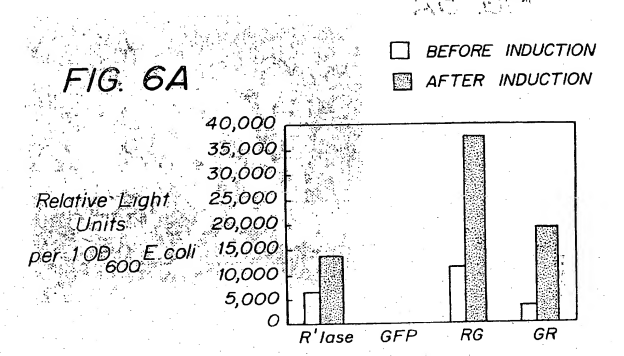
FIG. 58

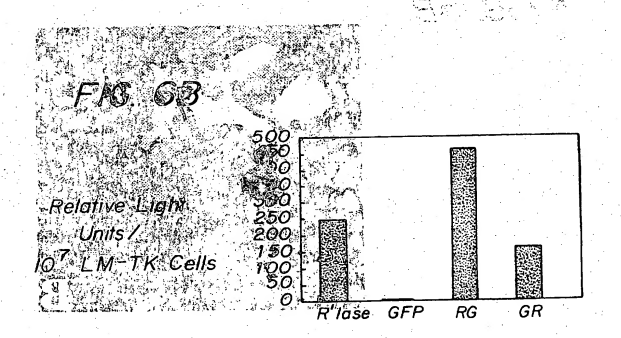
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FIG. 7

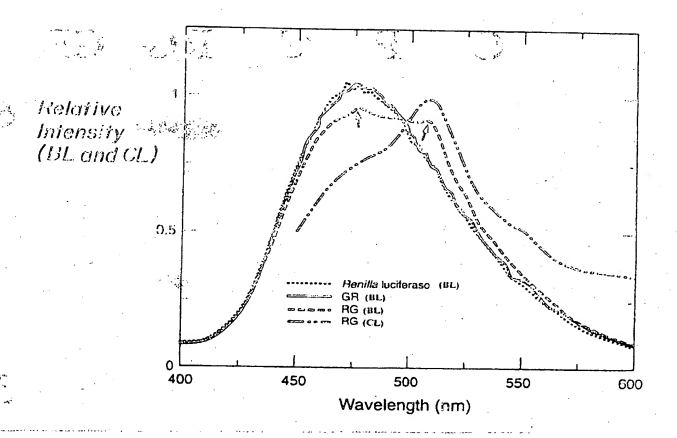
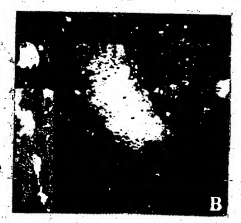


FIG. 9A

FIG. 9B



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F1G. 9D



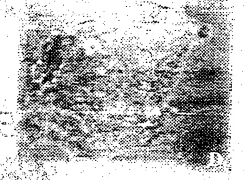
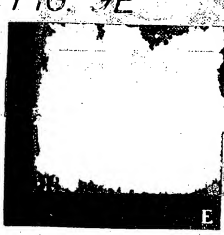


FIG. 3E

16. 9F





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FIG. 8

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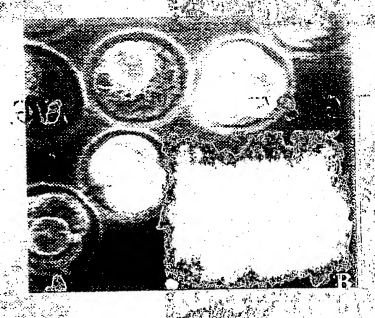
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FIG. MOA



FIG. 10B



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International application No.

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Electronic d	ata base consulted during the international search (name of data base and, where practicable	, search terms used)
	AT, EPOABS, JPOABS), STN (CAPLUS, BIOSIS) ms: luciforase, green fluorescent protein, renilla, aequorea, DNA, fusion, gene, antibody, mon	ocional
C. DOC	UMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y —	US 5,491,084 (CHALFIE et al) 13 February 1996, entire patent, especially column 1, lines 16-25 and claims	1,2, 6-9, 11, 13, 15-17, 19-21
A		3, 12, 14, 20
Y —	US 5,292,658 (CORMIER et al) 08 MARCH 1994, entire patent, especially claims.	1, 2, 6-9, 11, 13, 15-17, 19-21
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X Furth	er documents are listed in the continuation of Box C. See patent family annex.	
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Box 1 Observations where certain claims were	found unsearchable (Co	ontinuation of item-1-of-first sheet)
This international report has not been established in res	pect of certain claims unde	er Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not	required to be searched b	y this Authority, namely:
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This International Searching Authority found multiple	e inventions in this intern	national application, as follows:
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A. CLASSIFICATION OF SUBJECT MATTER: IPC (6):

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C12P 21/04, 21/06; C12N 1/20, 9/02, 15/09; C07K 14/00, 16/00; C07H 21/04

A. CLASSIFICATION OF SUBJECT MATTER: US CL :

435/69.1, 69.7, 189, 252.3, 320.1; 530/350, 388.1; 536/23.2, 23.4, 23.5

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING ar This ISA, found multiple inventions as follows: 2000 Productions (3) 2000 and 2000 and 2000 and 2000 and

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s)1-3, 6 and 7, drawn to a fusion protein having both luciferase and GFP activities.

Group II; claim(s) 4, 5 and 10, drawn to a monoclonal antibody against said fusion protein. 352 5

Group III, claim(s) 8, 9 and 11-17, drawn to a DNA encoding said fusion protein, a vector containing said DNA, a cell transformed with the same, a method-of-producing-said fusion-protein-using a transformed cell and lst-method of use of The first the region of the families maked all in the thereton inquiry moderal type of the state of the artistings of the first

Group IV, claim 18, drawn to a method of making a monoclonal antibody.

Group V, claim(s) 19-21, drawn to 2nd method of use of DNA encoding fusion protein.

The inventions listed as Groups I-V do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: a fusion protein of Group I, an antibody of Group II and a DNA of Group III are different compounds with different structures, functions and utilities. Luciforase and GFP as well DNAs encoding them and gene fusion constructs based on each of them are known in the prior art. An antibody against both proteins are known. Therefore, a fusion protein containing eithers. luciferase or GFP lacks a special technical feature with a DNA encoding thereof and an antibody against it.

Inventions of Groups IV and V are drawn to materially different methods. Method of Group IV employs immunization. of ah animal with a fusion protein and a hypridoma production, whereas a method of Group V employs a DNA construct encoding a fusion protein.

PCT Rule 1.475(d) does not provide for multiple products or methods within a single application and therefore, unity of invention is lacking with regard to Groups I-V. Anthor magneticities to the manufacture of the major of the first state of the stat

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